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- (22) Date of filing 2 Apr 1982
- (30) Priority data
- (31) 2327/81 2631/81 3278/81
- (32) 6 Apr 1981 22 Apr 1981 20 May 1981
- (33) Switzerland (CH)
- (43) Application published 1 Dec 1982
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- (52) Domestic classification A5B 822 826 M
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- (58) Field of search

- (54) Topical pharmaceutical compositions
- (57) A skin penetration pharmaceutical composition incorporating a difficultly skin-penetrable pharmacologically active agent, wherein the composition is in the form of a microemulsion formed from skin compatible excipients.

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The following corrections were allowed under Section 117 on 13 January 1984:

Front page, Heading (72), Inventor below Joachim Franz insert Jochen Ziegenmeyer

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- (58) Field of search A5B
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(54) Topical pharmaceutical compositions

(57) A skin penetration pharmaceutical composition incorporating a difficultly skin-penetrable pharmacologically active agent, wherein the composition is in the form of a microemulsion formed from skin compatible excipients.

Topical pharmac utical compositions

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|-----|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----|
| 5 | This invention relates to topical pharmaceutical compositions, particularly those containing pharmacologically active agents which only difficultly penetrate the skin horny layer. The therapeutic efficiency of a topical pharmaceutical composition depends upon inter alia the | 5 |
| | | |
| 10 | availability of the pharmacologically applied pharmacologically active agent can act at its site of active agent. Before any topically applied pharmacologically active agent can act at its site of action whether in the deeper dermal layers below the horny layer or elsewhere in the body it must penetrate the barrier of the horny layer of the skin (stratum corneum). The penetration of the stratum corneum is the rate-limiting step of the total percutaneous process and is accompanied by the creation of a reservoir of pharmacologically active agent, i.e. the deposition | 10 |
| 15 | of pharmacologically active agent on and in the layer. In the rare case the pharmacologically active agent is normally liquid and penetrates the skin layer efficiently, e.g. isosorbide dinitrate active agent is normally liquid and penetrates the skin layer efficiently, e.g. isosorbide dinitrate active agent is normally liquid and penetrates the skin layer efficiently, e.g. isosorbide dinitrate. | 15 |
| 20 | penetration of the pharmacologically active agent through the normy layer, especially for determining the specially administered in solid form. Often the pharmacologically active agent is capable of penetrating the skin horny layer when applied to the skin in a conventional system such as a triglyceride or paraffin ointment, but has a penetration flux of less than about 10 ⁻⁹ such as a triglyceride or paraffin ointment, but has a penetration flux of less than about 10 ⁻⁹ Mol cm ⁻² hour ⁻¹ . Such pharmacologically active agents are | 20 |
| 25 | One method to increase the penetration rate is to dissolve the skin-penetration pharmaceus, cally active agent in a non-toxic solvent which is skin compatible e.g. that does not cause skin irritation over an extended period of time as indicated in standard tests using human skin or more sensitive guineapig skin. The solutions may be applied in the form of macroemulsions, i.e. opaque oil-in-water or water-in-oil systems formed from water and water immiscible organic | 25 |
| 30 | solvents in the presence of an emulsitier. Such systems suffer from disadvantages especially in the case of difficultly skin-penetrable | 30 |
| • | The state of the state of the second state of | |
| | sition is in the form of a microemulation have particularly advantageous properties in respect of | |
| | difficultly skin-penetrable pharmacologically active agents. A recent review on microemulsions is by M. Rosoff p. 405 in Progress in Surface and | |
| 35 | A · 12 1070 Acception Proce A microphilishin is unificially recognition to the | 35 |
| • | the state of the sector of water-in-only pmilising wherein the dignifical of the particular | |
| | or droplets are less than about 1500 Angstrom units (150 nm) which is less than 1/4 of the wavelength of light. They do therefore not scatter visible light, the diameter of the particles or wavelength of light. | |
| | The state of the s | 40 |
| 40 | the state of the second of the second the second of the se | 40 |
| | isotropic or anisotropic. An anisotropic structure may however be observable using x-ray techniques. The particles in a microemulsion may be spherical but other structures are feasible, | |
| | ti it in a la collection de la collectio | |
| | the missions are produced from an emulsifier (a suffacially and a co-citiosino (i.o. | 4.5 |
| 45 | to the standard and distingt co-collibration in the internation tension between the | 45 |
| | oil-in-water phases to a very small amount (typically less than 1 dyne/cm). The microemulsions often form practically spontaneously and represent a single thermodynamically stable phase. In | |
| | and the macroamulations are thermodynamically unstable two phase systems, and in their | |
| | formation approximation the form of heating of fabig aditation is required. | 50 |
| 50 | the state of the s | 50 |
| | paints and foods. However, the formulation of microemulsions is to a certain extent largely empirical (see for example p. 34–56 in Microemulsions Theory and Practice, Ed. L. Prince, | |
| | A ORTHOLOGICAL TO A TO A TO A CONTROL PONCE OF THE PROPERTY OF | |
| | The state of a difficultive contraction by the contraction of the cont | 55 |
| 55 | from skin compatible excipients. J. Ziegenmeyer and C. Funrer in Acta Friannaceutical | 00 |
| | "" 10/ totrocyclin by droching and decand, hidyeyer, the composition is | |
| | the first transfer of the specific effect as the lettacount of the specific control of the specific co | |
| | pharmaceutical composition is too low. More importantly decanol is not skin compatible. For example in sensitive animal skin irritation tests, mod rate irritation of guinea pig skin and s vere example in sensitive animal skin irritation tests, mod rate irritation of guinea pig skin and s vere | 60 |
| 60 | | |
| | a in the state of the Editor Editor Editor (1947) of 140/, intercent the control of the control | |
| | Wiley, New York and Lond n. In I ss sensitive tests using human skin exposed to decanol v r a 24 h ur period, significant irritation has been observed, s e—for example p. 753 W. Kästner, a 24 h ur period, significant irritation has been observed. | |
| e i | a 24 h ur period, significant irritation has been observed, s each of example p. 756 been a 24 h ur period, significant irritation has been observed, s each of example p. 756 been a 24 h ur period, significant irritation has been observed, s each of example p. 756 been observed, s each observed p. 756 been observed, s each observed p. 756 been ob | 65 |
| ο: | 3. 300. Cosmon Chemistra (107 1) = 27 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | • |

suggested are not applicable for man. We have found that microemulsions may be made containing pharmacologically active agents and skin compatible excipients which show particularly advantag ous penetration properti s producing a penetration flux sufficient to produce a therapeutic effect in the deeper dermal 5 layers or through the systemic circulation as indicated in trials mentioned hereinafter. 5 In one aspect the present invention provides a skin penetration pharmaceutical composition incorporating a skin-penetrable pharmacologically active agent, wherein the composition is in the form of an microemulsion formed from skin compatible excipients. In another aspect the present invention provides a method of enhancing the penetration of a 10 skin-penetrable pharmacologically active agent through the skin which comprises applying the 10 active agent to the skin in the form of a microemulsion consisting of skin compatible excipients. In a further aspect the present invention provides the use of a microemulsion consisting of skin compatible excipients to administer percutaneously a skin-penetrable pharmacologically In yet a further aspect the present invention provides a process for the production of a skin-15 penetrable pharmaceutical composition which comprises forming a microemulsion from water and a skin-penetrable pharmacologically active agent and skin compatible excipients capable of functioning as a water-immiscible organic solvent, an emulsifier, and a co-emulsifier. The microemulsions may be produced in conventional manner for the preparation of topical 20 pharmaceutical compositions. The skin compatible pharmacologically active agent, water-20 immiscible organic solvent, water, emulsifier and co-emulsifier may be mixed, conveniently at a maximum of 100°C, e.g. from about 60° to about 95°C and the mixture is cooled. It is not important that a microemulsion be formed above 32°C. If a microemulsion is formed above 32°C then the phase inversions should preferably be 25 reversible. Indeed it is quite common that a milky macroemulsion may be formed at high 25 temperatures which on cooling passes through one or more cloudy transitional phases alternately with microemulsion phases. Desirably a microemulsion is produced throughout the temperature range of from about 20°C to about 32°C, preferably from about 15°C to about 35°C. The water-immiscible organic solvent may be for example a hydrocarbon or lipophilic ester. 30 An emulsifier is present to form an oil-in-water or water-in-oil emulsion wherein the oil is the water-immiscible organic solvent. The co-emulsifier contributes to the formation and the stability of the microemulsion. The chemical structure or chainlength of the co-emulsifier is a governing factor in controlling 35 the size of the droplets or particles in the emulsions and should match the structure or 35 chainlength of the hydrocarbon part of the emulsifier. The co-emulsifier should be compatible with the water-immiscible organic solvent forming the lipophilic phase. The organic solvent emulsifier and co-emulsifier should also be compatible with the pharmacologically active agent. Naturally it is possible that the same excipient acts as a water-immiscible organic solvent and 40 simultaneously as a co-emulsifier. Conveniently different excipients are used as organic solvent 40 and co-emulsifier, however. The microemulsions may be colourless or coloured, e.g. yellow. A suitable combination of an emulsifier with a co-emulsifier may be, for example, a watersoluble non-ionic emulsifier and a fatty alcohol of a suitable chain length. Another suitable combination may be a mixture of water-soluble and water-insoluble non-ionic tensides. Conveni-45 ently at least two of the water-immiscible organic solvent, co-emulsifier and emulsifier has a 45 chain length moiety of 12 to 20 carbon atoms. For any particular skin compatible pharmacologically active agent, water-immiscible organic solvent, water, emulsifier, and co-emulsifier system the relative amount of excipients can be varied and full phase equilibria diagrams may be drawn. It is sometimes more convenient merely 50 to obtain a microemulsion at any temperature, even above room temperature, from one set of 50 excipients in order to show they are compatible and then vary the amounts slightly to produce a suitable microemulsion at room temperature. As a very rough guide the microemulsion may contain:a) 0.01 to 15% of skin compatible skin-penetrable pharmacologically active agent, 55 b) 5 to 30%, e.g. 10 to 30%, of skin compatible water-immiscible organic solvent, 55 c) 10 to 30% of skin compatible emulsifier, d) 4 to 30% of skin compatible co-emulsifier, and e) 15 to 55% water. Where the same compound may act as, e.g. both water-immiscibl organic s Ivent and co-60 emulsifier, and in particular when anoth r co-emulsifier or organic s Ivent is omitted then a part 60 f the concentration of the compound (together with any other water-immiscible solvent present) may be reckoned as water-immiscibl solv nt and a part (together with any other co-emulsifier

pr sent) as co- mulsifier. Where the same excipient acts as both water-immiscible organic solv nt and co-emulsifier and the residue is no co-emuls and the residue is no co-emul

65 excipi nt may b present from 9 to 60% of the composition.

| The microemulsions of the invention may be in the control adults of the gals, which are semi-viscous, containing I se water. Some microgals may have appropriate viscoelastic properties to form swingling gets. In respect of any of the excipients mentioned hereinafter any aliphatic carboxylic acid may to straight-chain or branched and saturated or unsaturated, preferably with one or two double bonds. Any aliphatic alcohol is a univalent alcohol unless otherwise mentioned, and is preferal a secondary or especially a primary alcohol. They are branched or preferably straight-chain an are unsaturated with preferably one or two double bonds or especially saturated. Any glyceryl ether or ester is primarily etherified or esterified at one or both of the terminal glyceryl hydrox 10 groups. Suitable skin compatible excipients may be the following:— 1) an ester of an aliphatic (C ₀₋₁₂) alcohol with an aliphatic (C ₀₋₁₂) carboxylic acid, or 2) a hydrocarbon having a straight carbon (C ₁₂₋₃₂) chain substituted by from 5 to 16 methylgroups and having up to 6 double bonds, 15 may be suitable water-immiscible organic solvents. Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, isopropyl myristate and lauryl myristate. Examples of class 2) include terpenes such as polymethylbutanes and polymethylbutenes, e. 2, 6, 10, 15, 19, 23-hexamethyl-2,6, 10, 14, 18, 22 tetracosahexaene, also known as squalene (C ₀₋₁₄₋₃₂) and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from 3) a mone-sater of athylene glycol or propylene glycol with an aliphatic (C ₁₋₂₂) carboxylic acid. The suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of glycerol with an aliphatic (C ₁₋₂₂) carboxylic acid. The suitable for use as water-immiscible organic solvents and propylene glycol monomyr tata and preferably propylene glycol monolaurate. An example of class 3) is present as an organic solvent, then the scripien | | the form of liquids or preferably in the form of | |
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| In respect of any of the excipients mentioned hereinatter any aliphatic cardovy, automore 5 straight-chain or branched and saturated or unsaturated, preferably with one or two double bonds. Any aliphatic alcohol is a univalent alcohol unless otherwise mentioned, and is preferably are unsaturated with preferably one or two double bonds or especially sturated. Any glyceryl ether or ester is primarily etherified or esterified at one or both of the terminal glyceryl hydrox or ester is primarily etherified or esterified at one or both of the terminal glyceryl hydrox or ester is primarily etherified or esterified at one or both of the terminal glyceryl hydrox or surface and aliphatic (Ca ₃₋₁₈) alcohol with an aliphatic (Ci ₀₋₂₂) carboxylic acid, or 2) a hydrocarbon having a straight carbon (Ci ₁₃₋₂₃) chain substituted by from 6 to 16 methyl. groups and having up to 6 double bonds. 5 may be suitable water-immiscible organic solvents. Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, isopropyl myristate and lauryl myristate. Examples of class 1) include isopropyl laurate, hexyl laurate and isopropyl myristate, especially haxyl laurate. Examples of class 2) include terpenes such as polymethylbutanes and polymethylbutenes, 6 (C ₃ H ₂₀) and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C ₂₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/ or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the sc or different excipient may be present as a co-emulsifier. Some of this class and in aliphatic (Ci ₁₋₂₂) alcohol with an aliphatic (Ci ₃₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and propylene glycol monolaurate and propylene glycol monolaurate and propylene glycol monolaurate and propylene glycol monolaurate and p | | The microemulsions of the invention may be in the form of liquids or preferably in the form of gels, which are semi-viscous, containing I ss water. Some microgels may have appropriate | |
| 5 straight-chain or branched and saturated or unsaturated, preterally with other with sonds. Any aliphatic alcohol is a univalent alcohol unless otherwise mentioned, and is preferal a secondary or especially a primary alcohol. They are branched or preferably straight-chain an are unsaturated with preferably one or two double bonds or especially saturated. Any glycaryl ether or ester is primarily etherified or esterified at one or both of the terminal glycaryl hydrox 10 groups. Suitable skin compatible excipients may be the following:— 1) an ester of an aliphatic (C ₃₋₁₀) alcohol with an aliphatic (C ₁₀₋₂₂) carboxylic acid, or 2) a hydrocarbon having a straight carbon (C ₁₁₋₂₂) chain substituted by from 6 to 16 methyl-groups and having up to 6 double bonds, Tamples of class 1) include isopropyl leurate, hexyl laurate and decyl laurate, isopropyl myristate and lauryl myristate. Particularly suitable examples are isopropyl alurate, hexyl laurate and isopropyl myristate, especially hexyl laurate. Examples of class 2) include terpenes such as polymethylbutanes and polymethylbutanes, etc. 10, 15, 19, 23-hexamethyl-2, 6, 10, 14, 18, 22 tetracosahexaene, also known as squalane. (C ₃₋₃ H ₃₀) and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from Skin compatible excipients of glycerol with an aliphatic (C ₄₋₂₂) carboxylic acid, any be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the sc or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol group clause and propylene glycol monomyr tate and preferably propylene glycol monolourate. An example of class 5) is glyceryl caphylate. Examples of class 7) include double and specially d | | viscoelastic properties to form swinging gels. | |
| bonds. Any aliphatic alcohol is a univalent alcohol unites otherwise intertorious, and a phonose as ascondary or especially a primary alcohol. They are branched or preferably straight-chain an are unsaturated with preferably one or two double bonds or especially saturated. Any glycaryl ether or ester is primarily etheriford or esterified at one or both of the terminal glycaryl hydrox (10 groups. Suitable skin compatible excipients may be the following:— 1) an ester of an aliphatic (C ₂₋₁₀) alcohol with an aliphatic (C ₁₀₋₂₂) carboxylic acid, or 2) a hydrocarbon having a straight carbon (C ₁₁₋₂₂) chain substituted by from 6 to 16 methyl groups and having up to 6 double bonds. 5 may be suitable water-immiscible organic solvents. Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, isopropyl myristate and lauryl myristate. Examples of class 2) include terpenes such as polymethylbutanes and polymethylbutenes, e. 2, 6, 10, 15, 9,23-hexamethyl-2,6,10,14,18,22 tetracosahexane, also known as squalane. Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C ₆₋₂₂) carboxylic acid. The suitable of an aliphatic (C ₁₂₋₂₂) alcohol with lactic acid, or 5) a mono-or diester of glycerol with an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomory diester of any of classes 3), 4) or 5) is present as an organic solvent, then the sc or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 3) is myristyl or preferably lauryl lactace. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) and an aliphatic (C ₆₋₂₂) carboxylic acid. may be suitable for use as water-immiscible solvents or | 5 | attained about or branched and saturated or unsaturated, preterably with one or two occupion | 5 |
| a secondary or especially a primary alcohol. They are branched or pieceally saturated with preferably one or two double bonds or especially saturated. Any glyceryl ether or ester is primarily etherified or esterified at one or both of the terminal glyceryl hydrox 10 groups. Suitable skin compatible excipients may be the following:— 1) an ester of an aliphatic (C ₂₋₁₉) alcohol with an aliphatic (C ₁₀₋₂₂) carboxylic acid, or 2) a hydrocarbon having a straight carbon (C ₁₁₋₂₁) chain substituted by from 6 to 16 methylgroups and having up to 6 double bonds, groups and having up to 6 double bonds. Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, isopropyl myristate and lauryl myristate. Particularly suitable examples are isopropyl laurate, hexyl laurate and isopropyl myristate, especially hexyl laurate, and isopropyl myristate, especially hexyl laurate, and isopropyl myristate, especially hexyl laurate, isopropyl myristate, especially hexyl laurate, and isopropyl myristate, and preferably propylene glycol monolaurate. An example of class 3) include propylene glycol monolaurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 3) include propylene glycol monolaurate, and example of class 3) include propylene glycol monolaurate. An example of class 3) include propylene glycol monolaurate, and isopropylene glycol myristate, a | J | Landa Anti-cliphatic alcohol is a univalent alcohol unless otherwise mentioned, and is protected, | |
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| Suitable skin compatible excipients may be the following: 1) an ester of an aliphatic (C ₁₋₁₀) alcohol with an aliphatic (C ₁₀₋₂₂) carboxylic acid, or 2) a hydrocarbon having a straiglic tarbon (C ₁₋₂₃) chain substituted by from 6 to 16 methylgroups and having up to 6 double bonds, 15 may be suitable water-immiscible organic solvents. Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, isopropyl myristate and lauryl myristate. Particularly suitable examples are isopropyl laurate, hexyl laurate and isopropyl myristate, especially hexyl laurate. Examples of class 2) include terpenes such as polymethylbutanes and polymethylbutanes, especially hexyl laurate. Examples of class 2) include terpenes such as polymethylbutanes, and polymethylbutanes and polymethylbutanes, especially hexyl laurate. Examples of class 2) include terpenes such as polymethylbutanes, and polymethylbutanes and polymethylbutanes, especially hexyl laurate. Examples of class 2) include terpenes such as polymethylbutanes, and polymethylbutanes, and compared to the such as a polymethylbutanes, and polymethy | | are unsaturated with preferably one or two double bonds or especially saturated. Any gryssified at one or both of the terminal glyceryl hydroxy | |
| Suitable skin compatible excipients may be the following: 1) an ester of an aliphatic (C ₁₋₁₂) alcohol with an aliphatic (C ₁₋₁₂) carboxylic acid, or 2) a hydrocarbon having a straight carbon (C ₁₂₋₃₂) chain substituted by from 6 to 16 methylgroups and having up to 6 double bonds. To suitable water-immiscible organic solvents. Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, isopropyl myristate and lauryl myristate. Particularly suitable examples are isopropyl laurate, hexyl laurate and isopropyl myristate, especially hexyl laurate. Examples of class 2) include terpenes such as polymethylbutanes and polymethylbutenes, e. 2, 6, 10, 15, 19, 23-hexamethyl-2, 6, 10, 14, 18, 22 tetracosahexaene, also known as squalene. Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C ₆₋₂₂) carboxylic acid. May be suitable for use as water-immiscible organic solvents and /or co-emulsifiers. When a excipient of any of classes 3), 40 ro 5) is present as an organic solvent, then the sc or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably lauryl lactate. An example of class 5) is glycenyl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7) ethylene glycol glycerol ether having at least one free hydroxyl grouples as a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7) ethylene glycol glyceryl caprylate. Some of this class may be water-miscible when for example the polyethylene glycol monoisty has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7) thylene glycol glyceryl capted in USF 3, 288,824, the conter of which are hereby incorporated by reference, then the products may be water-immiscible as suitable for use as a water | 10 | | 10 |
| 1) an ester of an aliphatic (C ₃₋₁₈) alcohol with an aliphatic (L ₁₀₋₂₃) catrobyline do. 16 methylgroups and having up to 6 double bonds, 15 may be suitable water-immiscible organic solvents. Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, isopropyl myristate and lauryl myristate. Particularly suitable examples are isopropyl laurate, hexyl laurate and isopropyl myristate, especially hexyl laurate. Examples of class 2) include terpenes such as polymethylbutanes and polymethylbutenes, e.g., 6, 10, 15, 19, 23-hexamethyl-2, 6, 10, 14, 18, 22 tetracosahexaene, also known as squalene (C ₃₀ H ₃₀) and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C ₆₋₂₂) carboxylic acid. Any as ester of an aliphatic (C ₁₂₋₂₂) alcohol with lactic acid, or by a suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the scondifferent excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7) ethylene glycol glycerol ether having at least one free hydroxyl group and an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents, but they may suitable co-emulsifiers. Some of this class may be water-miscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable for use as an avater-immiscible organic solvent. Skin compatible excipients does not from 10 to 18, may be suitable for the analysis of cl | 10 | Suitable skin compatible excipients may be the following:— | |
| 2) a hydrocarbon having a straight carbon (c ₁₂₋₃₂) chain substitutes by from 5 to 15 metally groups and having up to 6 double bonds. Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, isopropyl myristate and lauryl myristate. Particularly suitable examples are isopropyl laurate, hexyl laurate and isopropyl myristate, especially hexyl laurate. Examples of class 2] include terpenes such as polymethylbutanes and polymethylbutanes, e Examples of class 2] include terpenes such as polymethylbutanes, and polymethylbutanes and polymethylbutanes, e (C ₆₀ H ₆₀) and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the sc or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol monolaurate and propylene glycol monomyr tate and preferably lauryl lactate. An example of class 5) is glycenyl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2–7)ethylene glycol glycerol ether having at least one free hydroxyl grot. Any skin compatible excipients chosen from 6) an ester of a poly(2–7)ethylene glycol glycerol ether having at least one free hydroxyl grot. Sand an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)ethylene glycol glycoryl cocoate. If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethylene glycol of molecular weight, on the suitable example is not pol | | 4) of an alimbatic (C) sicobol with an alignatic (C _{10,22}) carboxylic acid, or | |
| 15 may be suitable water-immiscible organic solvents. Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, proposed and suryl myristate. Particularly suitable examples are isopropyl aurate, hexyl laurate and isopropyl myristate, especially hexyl laurate. Examples of class 2) include terpenes such as polymethylbutanes and polymethylbutanes, etc., 10, 15, 19, 23-hexamethyl-2,6,10,14,18,22 tetracosahexaene, also known as squalane. (C ₃₀ H ₅₀) and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C ₄₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the sc or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol monolaurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl grou and inplatic (C ₄₋₂₂) carboxylic acid. may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)ethylene glycol glyceryl cocoate. If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethy en glycol of molecular weight 200 to 400, e.g. as described in USP 3,288,824, the conter of which are hereby incorporated by reference, then the products may be water-immiscible as usutable for use as an water-imm | | 2) a hydrocarbon having a straight carbon (C ₁₂₋₃₂) chain substituted by nom o to 70 metry. | |
| Examples of class 1) include isopropyl laurate, nexyl isurate and actyl myristate. Particularly suitable examples are isopropyl laurate, hexyl laurate and isopropyl myristate, especially hexyl laurate. 20 | 4 = | The authority material impriscible organic solvents. | 15 |
| myristate and lauryl myristate. Particularly suitable examples are isopropyl laurate, hexyl laurate and isopropyl myristate, especially hexyl laurate. 20 | 15 | Examples of class 1) include isopropyl laurate, hexyl laurate and decyl laurate, isopropyl | |
| Particularly suitable examples are isopropyl laurate, nexyl aurate and isopropyl myrister, especially hexyl laurate. 20 Examples of class 2) include terpenes such as polymethylbutanes and polymethylbutenes, et 2, 6, 10,15,19,23-hexamethyl-2,6,10,14,18,22 tetracosahexaene, also known as squalene. Claship, and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from 3) a mono-ester of sthylene glycol or propylene glycol with an aliphatic (C6,22) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the sc or different excipient may be present as a co-emulsifier. 30 Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably lauryl lactate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl grous and an aliphatic (C6,22) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7) ethylene glycol glyceryl cocoate. 40 If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethylene glycol of molecular weight 200 to 400, e.g. as described in USP 3,288,824, the content of which are hereby incorporated by reference, then the products may be water-immiscible as suitable for use as an water-immiscible organic solvent. Skin compatible excipient chosen from 7) aliphatic (C1,22) alcohol, or 8) an ester of an aliphatic (C6,22) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) i | | i | |
| Examples of class 2) include terpenes such as polymentryloutanes and polymentry. 2, 6, 10, 15, 19, 23-hexamethyl-2, 6, 10, 14, 18, 22 tetracosahexaene, also known as squalene (C₃₀H₅₀) and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C₆₋₂₂) carboxylic acid. 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the sc or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl grot and an aliphatic (C₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)ethylene glycol glyceryl coccate. If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethylene glycol of molecular weight 200 to 400, e.g. as described in USP 3,288,824, the conter of which are hereby incorporated by reference, then the products may be water-immiscible as suitable for use as an water-immiscible organic solvent. Skin compatible excipient chosen from 8) a nester of having at least one hydroxyl group of a poly-(2-10)-glycerol with an aliphatic (C₈₋₂₂) carboxylic acid, may be also s | | Particularly suitable examples are isopropyl laurate, nexyl laurate and isopropyl mynistate, | |
| 2, 6, 10,15,19,23-hexamethyl-2,6,10,14,18,22 tetracosanexaere, also known (C₃₀H₈₀) and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C₈₋₂₂) carboxylic acid, as ester of an aliphatic (C₁₂₋₂₂) alcohol with lactic acid, or 5) a mono-or diester of glycerol with an aliphatic (C₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the set or different excipient may be present as a co-emulsifier. 50 Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl ground an aliphatic (C₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)ethylene glycol glyceryl coccate. 40 If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethy ene glycol of molecular weight 200 to 400, e.g. as described in USP 3,288.824, the content of which are hereby incorporated by reference, then the products may be water-immiscible as suitable or use as an water-immiscible organic solvent. 5 Skin compatible excipient chosen from 7 an ester of having at least one hydroxyl group of a poly-{2-10}-glycerol with an aliphatic (C | | especially hexyl laurate. | 20 |
| (C ₃₀ H ₅₀) and the perhydro analogue, squalane. A particularly suitable example is squalane. Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C ₆₋₂₂) carboxylic acid, or 5) a mono-or disester of glycerol with an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the se or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl grounds and an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)ethylene glycol glyceryl cocate. 1 If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethylene glycol of molecular weight 200 to 400, e.g. as described in USP 3,288,824, the content of which are hereby incorporated by reference, then the products may be water-immiscible as suitable for use as an water-immiscible organic solvent. Skin compatible excipient chosen from 7 in the product of a poly-(2-10)-glycerol with an aliphatic (C ₆₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, having an HLB value of from 10 to 18, or 510 an ester of an aliphatic (C ₆₋₂₂) carboxylic acid with a) a polyethylene glycol sorbitan | 20 | a a 40 45 40 22 hovemothyl-2 6 10 14 18 22 tetracosanexaene, also known as squalene | |
| Skin compatible excipients chosen from 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (C ₆₋₂₂) carboxylic acid 4) an ester of an aliphatic (C ₁₋₂₋₂) alcohol with lactic acid, or 5) a mono-or diester of glycerol with an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the so or different excipient may be present as a co-emulsifier. So Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl grot and an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)thylene glycol glyceryl cocoate. If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethy ene glycol of molecular weight 200 to 400, e.g. as described in USP 3,288,824, the conter of which are hereby incorporated by reference, then the products may be water-immiscible as suitable for use as an water-immiscible organic solvent. Skin compatible excipient chosen from 7) aliphatic (C ₁₋₂₂) alcohol, or 8) an ester of having at least one hydroxyl group of a poly-{2-10}-glycerol with an aliphatic (C ₆₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2- octyl-decanol. Particularly suitable examples include tetradecanol and especially dodeca | | (Co.Hr.) and the perhydro analogue, squalane. A particularly suitable example is squalance. | |
| 4) an ester of an aliphatic (C₁₋₂₋₂) alcohol with lactic acid. or 5) a mono-or diester of glycerol with an aliphatic (C₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the se or different excipient may be present as a co-emulsifier. 30 Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl grounds and an aliphatic (C₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)ethylene glycol glyceryl cocoate. 40 If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethy ene glycol of molecular weight 200 to 400, e.g. as described in USP 3,288,824, the content of which are hereby incorporated by reference, then the products may be water-immiscible as suitable for use as an water-immiscible organic solvent. Skin compatible excipient chosen from 45 7) aliphatic (C₁₋₂₋₂) alcohol, or 8) an ester of having at least one hydroxyl group of a poly-(2-10)-glycerol with an aliphatic (C₂₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2-conditions of the products of an aliphatic (C₆₋₂₂) carboxylic acid with a) a polyethylene glycol on a set of an aliphatic (C₆₋₂₂) carboxylic acid with a) a pol | • | Clim and the avaining the chosen from | |
| may be suitable for use as water-immiscible organic solvents and/or co-emulsifiers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the se or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl ground and an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)ethylene glycol glyceryl cocoate. If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethy ene glycol of molecular weight 200 to 400, e.g. as described in USP 3,288,824, the conter of which are hereby incorporated by reference, then the products may be water-immiscible as suitable for use as an water-immiscible organic solvent. Skin compatible excipient chosen from 45 7) aliphatic (C ₁₂₋₂₂) alcohol, or 8) an ester of having at least one hydroxyl group of a poly-(2-10)-glycerol with an aliphatic (C ₂₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2-conditions of the site of the products of an aliphatic (C ₆₋₂₂) carboxylic acid with a) a polyethylene glycol sorbitan ether, 50 octyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C ₁₂ | | 3) a mono-ester of ethylene glycol or propylene glycol with an aliphatic (06-22) carboxy is used. | 25 |
| may be suitable for use as water-immiscible organic solvents and/ or Co-emitishers. When a excipient of any of classes 3), 4) or 5) is present as an organic solvent, then the se or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl grounds and an aliphatic (C ₆₋₂₂) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)ethylene glycol glyceryl cocoate. If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethy ene glycol of molecular weight 200 to 400, e.g. as described in USP 3,288,824, the contert of which are hereby incorporated by reference, then the products may be water-immiscible as suitable for use as an water-immiscible organic solvent. Skin compatible excipient chosen from 45 7) aliphatic (C ₁₂₋₂₂) alcohol, or 8) an ester of having at least one hydroxyl group of a poly-(2-10)-glycerol with an aliphatic (C ₆₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2-cotyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C ₁₂₋₁₈) alcohol, having an HLB valor of from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from | 25 | E) a manager diester of divideral with an alibhatic (C ₆₋₂₂) carboxyric acid, | |
| When a excipient of any of classes 3), 4) or 5) is present as an origanic solvent, then the se or different excipient may be present as a co-emulsifier. Examples of class 3) include propylene glycol mono-laurate and propylene glycol monomyr tate and preferably propylene glycol monolaurate. An example of class 4) is myristyl or preferably lauryl lactate. An example of class 5) is glyceryl caprylate. Any skin compatible excipients chosen from 6) an ester of a poly(2-7)ethylene glycol glycerol ether having at least one free hydroxyl ground an aliphatic (C ₈₋₂₂) carboxylic acid, may be suitable for use as water-immiscible solvents or co-emulsifiers. Some of this class may be water-miscible when for example the polyethylene glycol moiety has a higher molecular weight, and so will not be suitable as organic solvents, but they may suitable co-emulsifiers. An example is poly(7)ethylene glycol glyceryl cocoate. If the excipient is a transesterification product of a vegetable oil triglyceride and a polyethy ene glycol of molecular weight 200 to 400, e.g. as described in USP 3,288,824, the contert of which are hereby incorporated by reference, then the products may be water-immiscible as suitable for use as an water-immiscible organic solvent. Skin compatible excipient chosen from 37) aliphatic (C ₁₂₋₂₂) alcohol, or 38) an ester of having at least one hydroxyl group of a poly-(2-10)-glycerol with an aliphatic (C ₈₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2-cotyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 3) a mono-ether of a poly-ethylene-glycol with an aliphatic (C ₁₂₋₁₈) alcohol, having an HLB varian of from 10 to 18, or 50) an ester of an aliphatic (C ₆₋₂₂) carboxylic acid with a) a polyethylene glycol sorbitan ether, 60) the ester having an HLB valu of from 12 to 15 (HLB valu | | to suitable for use as water-immiscible organic solvents and/ or co-emulations. | |
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| ene glycol of molecular weight 200 to 400, e.g. as described in USP 3,286,824, the content of which are hereby incorporated by reference, then the products may be water-immiscible all suitable for use as an water-immiscible organic solvent. Skin compatible excipient chosen from 45 7) aliphatic (C ₁₂₋₂₂) alcohol, or 8) an ester of having at least one hydroxyl group of a poly-(2-10)-glycerol with an aliphatic (C ₆₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2-0 octyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C ₁₂₋₁₈) alcohol, having an HLB va of from 10 to 18, or 55 10) an ester of an aliphatic (C ₆₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indication of the hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | 40 | t also avaining to a transportarification product of a vegetable on triglycende and a polycury. | 40 |
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| Skin compatible excipient chosen from 7) aliphatic (C ₁₂₋₂₂) alcohol, or 8) an ester of having at least one hydroxyl group of a poly-(2-10)-glycerol with an aliphatic (C ₈₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2-octyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C ₁₂₋₁₈) alcohol, having an HLB va of from 10 to 18, or 55 10) an ester of an aliphatic (C ₈₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indication of the hydrophilic-lipophilic balance in an emulsifier and have been discussed extensively in a literature, see for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosmetic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | of which are hereby incorporated by reference, then the products may be water-immiscious and | |
| 7) aliphatic (C₁₂₋₂₂) alcohol, or 8) an ester of having at least one hydroxyl group of a poly-(2-10)-glycerol with an aliphatic (C₈₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2-octyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C₁₂₋₁₈) alcohol, having an HLB va of from 10 to 18, or 10) an ester of an aliphatic (C₆₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | Suitable for use as an water-immisciple organic soveric. | |
| 8) an ester of having at least one hydroxyl group of a poly-(2-10)-grycerol with an amphibite (C₈₋₂₂) carboxylic acid, may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2-octyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C₁₂₋₁₈) alcohol, having an HLB va of from 10 to 18, or 10) an ester of an aliphatic (C₈₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indication of the hydrophilic-lipophilic balance in an emulsifier and have been discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | 45 | : 7) alimbatia /C \ alcohol Or | 45 |
| may be also suitable co-emulsifiers. Examples of class 7) include dodecanol, tetradecanol, oleylalcohol, 2-hexyldecanol, and 2- 50 octyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C ₁₂₋₁₈) alcohol, having an HLB va of from 10 to 18, or 55 10) an ester of an aliphatic (C ₈₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | 8) an ester of having at least one hydroxyl group of a poly-(2-10)-glycerol with an allphane | |
| Examples of class 7) include dodecanol, tetradecanol, deviational, 2-hexyloecanol, and 2 octyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C ₁₂₋₁₈) alcohol, having an HLB va of from 10 to 18, or 55 10) an ester of an aliphatic (C ₈₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | (C ₆₋₂₂) carboxylic acid, | |
| 50 octyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. Preferably the alcohol is liquid at 32°C. Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C₁₂₋₁₈) alcohol, having an HLB va of from 10 to 18, or 55 10) an ester of an aliphatic (C₈₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | Evamples of class 7\ include dodecanol, tetradecanol, oleylarconol, 2-nexyluccanol, and 2 | |
| Skin compatible excipients chosen from 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C ₁₂₋₁₈) alcohol, having an HLB va of from 10 to 18, or 55 10) an ester of an aliphatic (C ₈₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | 50 | octyl-decanol. Particularly suitable examples include tetradecanol and especially dodecanol. | 50 |
| 9) a mono-ether of a poly-ethylene-glycol with an aliphatic (C₁₂₋₁₈) alcohol, having all HLB valof from 10 to 18, or 55 10) an ester of an aliphatic (C₈₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indication of the hydrophilic-lipophilic balance in an emulsifier and have been discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | Preferably the alcohol is liquid at 32°C. | |
| of from 10 to 18, or 55 10) an ester of an aliphatic (C ₈₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | Skin compatible exciplents chosen from | |
| 55 10) an ester of an aliphatic (C₈₋₂₂) carboxylic acid with a) a polyethylene glycol b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | of from 10 to 18, or | 55 |
| b) a saccharos c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | 5 | 5 10) an ester of an aliphatic (C ₆₋₂₂) carboxylic acid with | 55 |
| c) a sorbitan r d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | a) a polyethylene glycol | |
| d) a poly- thylene glycol sorbitan ether, 60 the ester having an HLB valu f from 10 to 18, may be suitable mulsifiers. Pref rably the emulsifiers hav an HLB valu of from 12 to 15 (HLB values ar an indicati of th hydrophilic-lipophilic balance in an emulsifier and have be n discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | | |
| 60 the ester having an HLB valu of from 10 to 18, may be suitable intuishers. Pref rably the emulsifiers have an HLB valu of from 12 to 15 (HLB values are an indication of the hydrophilic-lipophilic balance in an emulsifier and have been discussed extensively in literature, see for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosmitic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | d) a poly- thylene glycol sorbitan ether, | 60 |
| of th hydrophilic-lipophilic balance in an emulsifier and have be in discussed extensively in literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fiedler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | 60 | he ester having an HIR value of from 10 to 18, may be suitable invisitely. | |
| literature, s e for example Pharm.Act.Helv. (1969) 44, 9 and H.P.Fledler, Lexicon der Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1981, Editio Car | | of the hydrophilic-linophilic halance in an emulsifier and have be n discussed extensively in the | |
| Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1961, Editio Cal | | liameters a stor example Pharm Act Hely (1969) 44, 9 and H.P.Fledler, Lexicon del | |
| 65 AG. BRD). | _ | Hilfsstoff für Pharmazie, Kosm tic und angrenzende Gebiete, 2nd Edition, 1961, Editio Canton | 65 |
| 00 //2, 5//5/ | 6 | b AG, BRU). | |

| | , | |
|-----|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----|
| | A preferred exampl of class 9) is commercially available polyoxyethylene-(10)-ol yl ether. Preferably the microemulsions are made up from xcipients from class 1) and 2) as water-immiscible organic solv nts; especially class 1); class 7) as co-emulsifier and class 9) as emulsifier. | |
| 5 | The exact choice of organic solvent, emulsifier and co-emulsifier will depend on inter alia the | 5 |
| Ŭ | pharmacologically active agent used. | - |
| | The pharmacologically active agent may be any compound which can, penetrate the skin | |
| | horny layer, e.g. of molecular weight up to about 3,000, although higher molecular weight | |
| | compounds may possibly be used. | |
| 10 | In general the molecular weight of the pharmacologically active agent is conveniently below | 10 |
| | 1000. Conveniently the active agent has a good hydrophilic/lipophilic balance. The molecule of the active agent for example may be conveniently structurally compact, may contain aromatic | |
| | groups and conveniently does not contain many reactive groups such as hydroxyl groups. | |
| | The microemulsions of the invention are capable of containing very high amounts of active | |
| 15 | agents, e.g. from 5% up to 15% or even up to 20% of the total weight. When a systemic | 15 |
| | action is desired, the pharmacologically active agent should be sufficiently active to be able to | |
| | produce a systemic therapeutic effect when penetration the skin at rate of the order of 10 ⁻⁸ | |
| | Mole cm ⁻² hour ⁻¹ . When a local action in the deeper dermal layer is required, then a skin | |
| | penetration flux of 10 ⁻⁹ Mole cm ⁻² hour ⁻¹ may be sufficient. Suitable agents may be for | |
| 20 | example those with an, e.g. oral, daily dose of about 0.1 to about 20 mg, preferably up to 1 | 20 |
| | mg. The microemulsions of the invention may be indicated for the systemic administration of any | |
| | active agent. They may be conveniently used for prophylactic agents and myotonolytics. The | |
| | microemulsions of the invention may be indicated for the administration of pharmacologically | |
| 25 | active agents which act under the horny layer, e.g. anti-acne agents and anti-fungal agents. | 25 |
| | Examples of active agents include | |
| | (E)-N-methyl-6,6-dimethyl-N-(naphthylmethyl)hept-2-en-4-inyl-1-amine, proquazone, | |
| | (E)-N-methyl-N-(1-naphthylmethyl)-3-phenyl-propen-2-yl-amine (hereinafter naftifin), | |
| 30 | 4-(1-methyl-4-piperidylidene)-4H-benzo[4,5]cyclohepta-[1,2-b]thiophen-10(9H)-one (hereinafter ketotifen), | 2Ò |
| 30 | 4-(1-methyl-4-piperidylidene)-9,10-dihydro-4H-benzo-(4,5)-cyclohepta(1,2-b)-thiophene (herei- | 30 |
| | nafter pizotifen), griseofulvin, fluocinolone acetonide, Triamcinolone acetonide, and 14-0-[5-(2- | |
| | amino-1,3,4-triazol-yl)thioacetyl]-dihydro-mutiline, and preferably | |
| | (+)-1-methyl-2-[2-(α-methyl-p-chlorodiphenyl-methoxy)-ethyl]-pyrrolidine (hereinafter clemastine) | |
| 35 | and especially 5-chloro-4-(2-imidazolin-2-ylamino)-2,1,3-benzothiadiazole (hereinafter tizanidine). | 35 |
| | In respect of clemastine a microemulsion preferably contains any of the following concentra- | |
| | tions:— 5 to 15% of clemastine. | |
| | 5 to 30% of an water-immiscible organic solvent. | |
| 40 | 15 to 25% of an emulsifier. | 40 |
| | 5 to 25% of a co-emulsifier. | 70 |
| | 10 to 45% of water. | |
| | More preferably a microemulsion contains any of the following concentrations:— | |
| A E | 7.5 to 12.5% of clemastine. | |
| 45 | 7.5 to 28.5% of water-immiscible organic solvent. 19.5 to 22% of an emulsifier. | 45 |
| | 7.5 to 22.5% of a co-emulsifier. | |
| | 13 to 42% of water | |
| | More especially a clemastine microemulsion contains any of the following concentrations:— | |
| 50 | 8 to 12% of clemastine. | 50 |
| | 8 to 27% of water-immiscible organic solvent. | |
| | 20 to 21% of an emulsifier. | |
| | 8 to 21% of a co-emulsifier. 15 to 40% of water. | |
| 55 | The excipients are preferably chosen from class (1) as defined above, as organic solvent. | EE |
| 00 | The excipients of class (3) as defined above may be present as organic solvent or co- | 55 |
| | emulsifier, especially propylene glycol mono-laurate. The co-emulsifier alternatively is an | |
| | excipient of class (6) as defined above especially poly(7)ethylene glycol glyceryl cocoate, or | |
| | propylene glycol myristate. The preferr d emulsifi r is chosen fr m class (9) as defined ab ve, | |
| 60 | especially polyoxyethylene (10) oleyl eth r e.g. having an HLB valu f about 12 to 13. | 60 |
| | With clemastine microgels containing high concentrations of cl mastine can be produced | |
| | whereas it is very difficult to pr duce stable macroemulsions containing such high clemastin concentrations. | |
| | In the respect of tizanidine a microemulsion preferably contains any of the following | |
| 65 | concentrations:— | 65 |
| | | |

| | 6 to 10% of tizanidine. 15 to 25% of water-immiscible organic solvent. | |
|----|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----|
| | 15 to 25% of water-infrastric organic sorroin. 15 to 25% of an mulsifier. | |
| | 5 to 10% of a co-emulsifier. | 5 |
| 5 | 30 to 35% of water. Preferably the microemulsion contains any of the following concentrations:— | |
| | 7.5 to 8.5% of tizanidine. | |
| | 19.5 to 21.5% of water-immiscible organic solvent. | |
| | 19 to 22% of an emulsifier. | 10 |
| 10 | 5.5 to 21.5% of a co-emulsifier. | 10 |
| | 32 to 42% of water. More particularly the microemulsion contains any of the following concentrations:—— | |
| | 8 to 8.4% of tizanidine. | |
| | 20 to 21% of water-immiscible organic solvent. | 4- |
| 15 | 20 to 21% of an emulsifier. | 15 |
| | 6.2 to 8.4% of a co-emulsifier. | |
| | 33 to 42% of water. Naturally the choice of water-immiscible organic solvent, emulsifier and co-emulsifier for a | |
| | tom pharmacologically active adent to pharmacologically active | |
| 20 | agent, and in some cases a particular excipient may be suitable in one system as e.g. an water | 20 |
| 20 | · · · · · · · · · · · · · · · · · · · | |
| | The all of the pharmacoutical composition may be adjusted to a skin companior pri with | |
| | appropriate acids or bases, preferably weak acids or bases e.g. lactic or acetic acid. It is preferred that the pharmacologically active agent is at least partially present in free form, e.g. | |
| 25 | free base form as the skin penetration may be increased. Conveniently the pH of the | 25 |
| 25 | -i | |
| | Other skin compatible agents may be present, e.g. Water-Misciple solvents such as propylene | |
| | the lead asked and isopropagal or water soluble tilm-torming agents used in cosmetic | |
| | preparations, e.g. partially hydrolysed collagen yielding medium-weight polypeptides, to diminish solvent evaporation after rubbing on the skin. | 30 |
| 30 | At a collection chould be composed of components that die skin companies | |
| | The components should be non-toxic, non-allergic and well-tolerated by the skill tissue. Buth | |
| | and he chosen by standard acute and coronic tests. | |
| | The tests may be effected on human skin or with more sensitive animal skin, e.g. guinea-pig | 35 |
| 35 | skin. The microemulsions of the invention are indicated for use in the percutaneous administration | |
| | the skin benefit and the agents because of the skin benefitation children and the | |
| | and the microamulations to contain large amounts of pitallideologically delive agonts. | |
| | The skin-penetration enhancing effect may be observed in standard in vito and in vivo tests | 40 |
| 40 | for example using human skin. One in vitro test is the well-known diffusion test which may be effected according to the | |
| | win single described by U. Schaeffer et al in Adv. Pharmacol, Iner. (Proc. / III Int. Cong.) natification. | |
| | A AAA AAF /4A7A) -J L. V FABAA PARASMAN LIXIAM LIXIAM LIXIAM II. JUHGUUD CEU PPE CO CE | |
| | in Current Problems in Dermatology / Ed. L.A. SIMON et al., Natyel, basel (1970), and sixti | 45 |
| 45 | Franz et al, Arch. Dermatol. Res (1981), 271:275–282, using isolated human skin. Microemulsions with the pharmacologically active agent in radio-active labelled form are | |
| | and the indicated pieces of unbroken human abdominal skin of about 2 square certificates in | |
| | at an amount of about 5 to about 10 mg of microemusion per square continuous. The | |
| | This is maintained at 20°C as a harrier between an upper champer and physiological sailine | 50 |
| 50 | placed in a lower chamber. After 100,300 and 1000 minutes at 32°C the skin is fixed on a stopper. The residue is removed from the skin surface by a cotton swab and the radioactivity | •- |
| | The bear lever in removed by stripping and the radioactivity is usualliniou in oddin | |
| | individual etripping. The remaining skin is congedied and sliced life sections of about 20 10 m | |
| | with a microtome. The radioactivity in the various slices is determined. The radioactivity in | 55 |
| 55 | aqueous saline in contact with the underside of the skin is also measured. Since the penetration of the pharmacologically active agent through the horny layer represents | |
| | in annual the rate limiting eten the amount of pharmacologically active agent that has pade a | |
| | the beautifued in relations to the ever mic activity. This traction of prigning coloring con- | |
| | | 60 |
| 60 | thick), lower corium (ca 1000 microns thick) and sub-cutis (ca 1500 microns thick), w uld in vivo be remov d by the capillary system into the blood stream and hence into the general | 30 |
| | a landada m | |
| | The province the fraction of the pharmacologically active agent that has pinetrated the | |
| | L revelop r offer 16 hours and is present in the deeper dermal layers is intensition to give a | e e |
| 6 | 5 mean percutaneous genetration flux (F) on the basis of a number of trials (n) as well as a | 65 |

25

percutaneous resorption quota in % of th appli d dose (RQ). Results obtained are as follows:—

| 5 | Example* No | F(0-16 hours)X 10 ⁸ Mol cm ⁻² hour ⁻¹ | n | RQ (%) |
|----|-------------|---------------------------------------------------------------------------|----|--------|
| | 1 | 2.6 ± 0.5 | 8 | 24% |
| | 3 | 1.4 ± 0.3 | 20 | 14% |
| 10 | 4 | ca 1.6 | 4 | 13% |
| | 5 | ca 2.6 | 4 | 21% |
| | 13 | 1.3 ± 0.01 | 12 | 12% |
| | 14 | 1.7 ± 0.7 | 8 | 12% |
| | 17 | ca 1.2 | 4 | 15% |
| 15 | 18 | ca 1.7 | 4 | 25% |
| | 20 | ca 1.3 | 4 | 12% |
| | 25 | 1.6 ± 0.6 | 8 | 13% |

*The examples are listed hereinafter.

20

In vivo trials may be effected, e.g. including a comparative oral and percutaneous administration of the pharmacologically active agent in a cross-over study in a healthy subject.

In one study 480 mg of a microemulsion in the form of a gel as described in Example 1 containing 40 mg of active agent, tizanidine, was applied behind the ear, or a tablet containing 25 4 mg tizanidine, was administered orally.

The urine was collected over 72 hours and the amount of unchanged active agent and corresponding two metabolites were measured separately.

The results obtained were as follows:-

| 30 ——— | | |
|-----------------------------|--------------------------------------------------------|----------------------------------------------------------|
| Period after administration | unchanged drug after oral administration [µg/hr] | unchanged drug after percutaneous administration [µg/hr] |
| 35 0-2 | 3.08 | 0.03 |
| 2-4 | 1.61 | 1.01 |
| 4–6 | 0.53 | 1.81 |
| 6-8 | 0.24 | 1.33 |
| 8-12 | 0.04 | 3.36 |
| 40 12-24 | | 4.16 |
| 24-36 | | 2.54 |
| 36-48 | | 1.57 |
| 48-60 | _ | 1.10 |
| 60-72 | | 1.07 |
| 45 | | |
| Cumulative % | | • |
| absorption | oral | percutaneous |
| of tizanidine | 0.28% | 0.37% |
| of Metabolite A | 2.5 % | 0.4 % |
| 50 of Metabolite B | 1.1 % | 0.16% |
| | | |

The above results confirm the significant percutaneous absorption obtained in the in vitro tests, and indicate a sustained-release effect. Additionally the relatively lower amount of metabolite found indicates a significantly lower first pass effect.

100 mg of the clemastine composition of Example 3 (containing 10 mg clemastine) is applied behind the ear of 2 or 3 subjects (age 18 to 38 years) corresponding to an amount of active agent of 10 mg f clemastine.

The amount of active agent in the urine is determined acc rding to the principles of 60 R.Tham.Arzn im.Forsch. (1978) 28 (1), 1017.

60

55

| 5 | Period after administration hours | active ag nt in urine [µg/hr] | Subjects | | |
|----|----------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------|-------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---|
| 10 | 0-6 6-8 8-12 12-24 24-36 | -0.486 ± 0.164 0.890 ± 0.384 1.042 ± 0.621 1.101 ± 0.422 | 3 3 3 3 | | 1 |
| 15 | 36-48 48-60 60-72 | 1.469 ± 0.455 0.504 ± 0.211 0.231 ± 0.05 | 3 2 2 | | 1 |
| | ar o tubo | elimination of unch | nanged dru | g 0.664 ± 0.183 | |
| 20 | In comparison unchanged drug The results sho 36 hours after a | 2 mg of clemasting in the urine. | e given ora | lly over 72 hours yield 7.10% ± 0.46% of the maximum concentration in the urine occurring quota of about 10% of the clemastine topically | 2 |
| 25 | systemic action of that topical adm. The present in tizaniding as actions. | of the pharmacolog inistration of tizanid evention according ive agent. In anothe | ically active line is feas provides a er aspect th | re present invention provides a method of topically | |
| 30 | The penetration hour 1 to produce local ac | ice a systemic action | ay thus be n and in th dermal laye | e order of about 1 × 10 ⁻⁹ Mole cm ⁻² hour ⁻¹ to rs and the concentration of pharmacologically | |
| 35 | The amount of present invention agent observed skin area treated | n will depend inter in the in vitro or in depend or in depend on the microements as suitable daily | active age alia on the vivo tests, ulsion, the | penetration rate of the pharmacologically active the potency of the active agent, the size of the part of the body treated and the duration of action out from 5 to 20 times the dose effective in oral | |
| 40 | administration, and a general as a microemulsions. In the case of | and the dose may builtable application of the invention mail liquid preparations | area is from ay be appli to the micro | d if longer duration than 1 day is required. In about 1 to about 40 square centimeters. The ed in conventional manner. It is emulsion can be applied for example from a oper arm, or from a plaster soaked with the the case of semi-solid microgels these may be | |
| 45 | rubbed in the sl For example i mg, and this ma | kin. in the case of tizani ay last for up to 3 c | dine and cl days. The n | emastine a suitable single dose is from 10 to 50 nicroemulsions of the invention may be used for tarmaceutically active agents are used for, e.g. anidine as myotonolytic, anti-depressant or minor | |
| 50 | tranquillizer. The microem active agent when | ulsions of the inven | tion may e in the hore | nhance the penetration of the pharmacologically by layer of the epidermis. A depot effect may then a agent slowly passes into the systemic circulation longlasting concentration of active agent in the | |
| 5 | blood (retard ef characterized by | fect). The blood cory the absence of an | ncentration initial drug | achieved by percutaneous delivery may be concentration blood peak in contrast to oral Additionally the accumulated pharmacologically local eff ct if the pharmacologically active agent is | |
| 60 | locally activ The microem macroemulsion | s. For example they | ntion may in ge | n general poss ss significant other advantages ver neral be thermodynamically stable, and show little | |
| 6 | properties n the easily rubbed in if desired. The | he skin surface. The | ey don't in little gr asy jnificantly (| general stick t the surfac f the skin but may be feeling behind and may be washed ff with water lehydrated as the single water-containing phase | |

| 5 | availabl from Atlas, Essen, or VOLPO 10 having an HL Polyethylene glycol glycerol from Gatte-fosse, Boulogne, Hexyllaurate is for example | ether is f r example eith r BRIJ 97 having an HLB value of 12.4 W. G rmany, B valu of 12.4 available from Cr da, Humb rside, UK. fatty acid ester is for example brand Labrafil M 1944 S available, France. brand CETIOL A, available from Henkel, Düsseldorf. | 5 |
|----|-------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----|
| 10 | Lactic acid is a 90% pure a collagen-derived cosmetic n Company, Northfield, III, US | | 10 |
| 15 | Ceraphyl 50 from Van Dyk, Further details on these p Pharmazie, Kosmetik und a | brand Ceraphyl 31, and myristyl lactate is for example brand Belleville, N.J., USA. broducts can be obtained from Fiedler H.P. Lexikon der Hilfsstoffe für ingrenzende Chemie, 2nd Edition, Editor Cantor, the contents of ted by reference, or their suppliers. | 15 |
| 20 | EXAMPLE ONE: Tizanidine 500 g of a mixture having sition:— | | 20 |
| | | Per cent | |
| 25 | Fizanidine Isopropyl laurate Polyoxyethylene (10) | 8.2 20.5 | 25 |
| | oleyl ether | 20.5 (Brij 97) | |
| | Dodecanol | 6.5 | |
| 30 | Water Lactic acid . | 41.0 3.3 | 30 |
| 35 | temperature by cooling the | water bath at about 90°C. The mixture is allowed to cool to room water 1°C per minute rious different phases are observed as follows:—— | 35 |
| | Phase | Temperature | |
| 40 | Milky macro-emulsion Transitional light cloudy phase | 92-72°C 72-70°C | 40 |
| 45 | Microemulsion transpa- rent phase Transitional light cloudy phase | 70-66°C 66-63°C | 45 |
| | Microemulsion transparent phase | 63-51°C | |
| 50 | Transitional light cloudy | | |
| 50 | phase Microemulsion transparent | 51-46°C | 50 |
| | phase | 46°—room temperature | |

The cooled gel is filled into metal tubes.

| Active age Example ingredient | Activ ingre | Active agent ingredient | Org. Solvent | | Co-Emulsifier | | Emulsifier | | dist. water | Additional excipients | 1 |
|----------------------------------|----------------|----------------------------|------------------------------------|-------|------------------------|------|------------------------------------------|-------|----------------|----------------------------------|------|
| | No. | % | В | | q | 8 | v | % | | % | .9 |
| 2 | - | 1% | Hexyllaurate | 23% | Poly(7)ethylene-glycol | 26% | Polyoxyethylene- 10-oleyl ether | 20% | 29.7% | 29.7% anhydrous acetic acid | 0.3% |
| ო | 7 | 10% | Hexyllaurate | 10% | Poly(7)ethylene-glycol | 20% | Polyoxyethylene- 10-oleyl ether *A | 20% | 38.5% | 38.5% anhydrous acetic acid | 1.5% |
| 4 | ო | 8.2% | 2,6,10,15,19,23- Hexamethyl-te- | 20.5% | Dodecanol | 6.5% | Polyoxyethylene- 10-oleyl ether | 20.5% | 40.6% | 40.6% lactic acid 90% | 3.7% |
| വ | က | 8.2% | Isopropylmyri- state | 20.5% | Dodecanol | 6.5% | Polyoxyethylene- 10-oleyl ether | 20.5% | 40.6% | 40.6% lactic acid 90% | 3.7% |
| 9 | က | 8.2% | Isopropylmyri- state | 20.5% | 20.5% Tetradecanol | 6.5% | Polyoxyethylene- 10-oleyl ether | 20.5% | 34.6% | 34.6% lactic acid 90%, Colla- | 3.7% |
| 7 | 4 | 8.3% | Isopropyl- laurate | 20.5% | Dodecanol | 6.5% | Polyoxyethylene- 10-oley ether | 20.5% | 41.3% | 41.3% lactic acid 90% | 2.9% |

| active in- Example gredient* | activ gred | active in- gredient* | Org. solvent | | Co-emulsifier | | Emulsifier | | dist. Additional water excipients | |
|---------------------------------|---------------|-------------------------|------------------------------------------------|-------|---------------|------|------------------------------------|-------|--------------------------------------|-------|
| | è. | % | o | | þ | % | · U | % | | % |
| ω | 5 | 8.3% | Isopropyl Iaurate | 20.6% | Dodecanol | 89.9 | Polyoxyethlyene- 10-oleyl ether | 20.6% | 41.2% lactic acid acid 90% | 2.7% |
| 6 | - | 1.0% | Isopropy aurate | 20.0% | Dodecanol | 7.0% | Polyoxyethylene- 10-oleyl ether | 18.0% | 53.7% lactic acid acid 90% | 0.3% |
| 10 | 9 | 0.5% | 2,6,10,15,19, 23-Hexamethyl- tetracosane | 21.0% | Dodecanol | 6.5% | Polyoxyethylene- 10-oleyl ether | 21.0% | 26.0% Polyethyl- englycol 400 | 25.0% |
| Ξ. | 7 | 0.2% | 2,6,10,15,19, 23-Hexamethyl- tetracosane | 20.5% | Dodecanol | 8.8% | Polyoxyethylene- 10-oelyl ether | 20.5% | 80.09 | |
| 12 | ω | 0.1% | sopropy aurate | 22.5% | Dodecanol | 8.0% | Polyoxyethylene- 10-oleyl ether | 22.5% | 46.9% | |
| 13 | ო | 8.2% | 2,6,10,15,19, 23-Hexamethyl- tetracosene | 20.5% | Dodecanol | 6.5% | Polyoxyethylene- 10-oleyl ether | 20.5% | 41.0% lactic acid 90% | 3.3% |
| 4 | ო | 8.2% | Isopropyl laurate | 20.5% | Dodecanol | 6.5% | Polyoxyethylene- 10-oleyl ether | 20.5% | 41.0% lactice acid 90% | 3.3% |

| active in- Example gredient | activ gredi | active in- gredient | Org. solvent | | Co-emulsifier | | Emulsifier | | dist. water | Additional excipients | % |
|--------------------------------|----------------|------------------------|------------------------------------------------|-------|------------------------------------------------|-------|------------------------------------|-------|----------------|--------------------------|------|
| | No. % | 8 | ro. | | S | 8 | U | % | | | % |
| 22 | 2 | 10% | Propylene gly- col mono-laurate | 13% | Poly(7)ethyl- 26% ene-glycol-gly- cerylcocoate | 26% | Polyoxyethylene- 10-oleyl ether | 20% | 31% | | |
| 23 | 8 | 10% | Propylene gly- col mono-laurate | 13% | | . 76% | Polyoxyethylene- 10-oleyl ether | 20% | 16% | Alcohol (96%) | 15% |
| 24 | က | 8.2% | lsopropyl myristate | 20.5% | Dodecanol | % | Polyoxyethylene- 10-oleyl ether | 20.5% | 39.1% | 39.1% lactic acid 90% | 3.7% |
| 25 | ო | 8.2% | 2,6,10,15,19, 23-hexamethyl- tetracosane | 20.5% | Dodecanol | 6.5% | Polyoxyethylene- 10-oleyl ether | 20.5% | 41% | lactic acid 90% | 3.3% |

| 5 | *Table of pharmacologically active agents 1. (E)-N-m thyl-N-(1-naphthylmethyl)-3-phenyl-prop n-2-ylamine. 2. (+)-1-methyl-2-[2-(α-methyl-p-chlorodiphenyl-methoxy)-ethyl]-pyrrolidin 3. 5-chloro-4-(2-imidazolin-2-ylamino)-2,1,3-benzothia-diazole. 4. 4-(1-methyl-4-piperidylidene)-4H-benzo[4,5]cyclohepta[1,2-b]thiophen-10(9H)-one. 5. 4-(1-methyl-4-piperidylidene)-9,10-dihydro-4H-benzo-(4,5)-cyclohepta-(1,2-b)-thiophene. 6. Griseofulvin. | 5 |
|-----|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----|
| 10 | 7. Fluocinolone acetonide. 8. Triamcinolone acetonide. 9. 14-0-[5-(2-amino-1,3,4-triazolyl)thioacetyl]-dihydro-mutiline, also known as 14-[5-amino-4H-1,2,4-triazol-3-yl)-thio-acetoxy]-14-deoxy-19,20-dihydromutilin. | 10 |
| 15 | **Table of commercial products A BRIJ 97 HLB value 12.4 (ATLAS) B VOLPO 10 HLB value 12.4 (CRODA) C CETIOL HE (HENKEL) D LAFABRIL 19445 (GATTEFOSSE) | 15 |
| 20 | Colladerm 350: A solution of a Zn salt of a highly purified cosmetic polypeptide of collagen (STEPHAN CHEMICAL COMPANY). | 20 |
| | CLAIMS 1. A skin penetration pharmaceutical composition incorporating a skin-penetrable pharmacologically active agent, wherein the composition is in the form of a microemulsion formed from | • |
| 25 | skin compatible excipients. 2. A composition as claimed in claim 1 wherein the composition is in the form of a microgel. | 25 |
| 30 | 3. A composition as claimed in claim 1 or 2 wherein the active agent is a difficultly skin-penetrable active agent. 4. A composition as claimed in claim 3 comprising from 5 to 30% by weight of a water-immiscible skin compatible solvent. | 30 |
| | 5. A composition as claimed in any preceeding claim containing from 4 to 30% by weight of a skin compatible emulsifier. 6. A composition as claimed in any preceeding claim comprising 10 to 30% by weight of a | |
| 35 | skin compatible co-emulsifier. | 35 |
| 40 | 8. A composition as claimed in any preceding claim containing 0.01 to 15% by weight of skin-penetrable pharmacologically active agent. 9. A composition as claimed in claim 8 containing from 5 to 15% by weight of skin-penetrable pharmacologically active agent. 10. A composition as claimed in any preceding claim containing a skin compatible ester of | 40 |
| 4.5 | an aliphatic (C ₃₋₁₈) alcohol with an aliphatic (C ₁₀₋₂₂) carboxylic acid. 11. A composition as claimed in claim 10 wherein the ester is chosen from isopropyl laurate, hexyl laurate, decyl laurate, isopropyl myristate and lauryl myristate. | 45 |
| 45 | 12. A composition as claimed in claim 10 wherein the ester is isopropyl laurate, hexyl laurate or isopropyl myristate. 13. A composition as claimed in claim 10 wherein the ester is hexyl laurate. | |
| 50 | 14. A composition as claimed in any preceding claim containing a skin compatible hydrocarbon having a straight carbon (C ₁₂₋₃₂) chain substituted by from 6 to 16 methyl groups and having up to 6 double bonds. | 50 |
| | 15. A composition as claimed in claim 14 containing squalane. 16. A composition as claimed in any preceding claim containing a skin compatible monoester of ethylene glycol or propylene glycol with an aliphatic (C ₆₋₂₂) carboxylic acid. | |
| 55 | 17. A composition as claimed in claim 16 wherein the ester is propylene glycol monolaurate or propylene glycol monomyristate. 18. A composition as claimed in any prece ding claim where in the ester is a skin compatible. | 55 |
| | ester f an aliphatic (C ₁₂₋₂₂) alcohol with lactic acid. 19. A composition as claim d in claim 18 wherein the ester is myristyl lactate or lauryl lactat. | |
| 60 | 20. A c mp sition as claimed in any prec ding claim containing a skin compatible aliphatic | 60 |
| | 21. A composition as claimed in claim 20 wherein the alc hol is d d can I, t tradecanol, oleyl alcoh I, 2-hexyldecanol r 2-octyldecanol. 22. A comp sition as claimed in claim 20 wherein the alcohol is dodecanol. | |
| 65 | | 65 |

| | a poly(2–7)ethylene glycol glyc rol ther having at least one fre hydroxyl group and an aliphatic (C_{6-22}) carboxylic acid. | |
|----|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----|
| | 24. A composition as claim d in claim 23 wherein the ester is p ly(7)ethylene glyc I glyceryl c coate. | |
| 5 | 25. A composition as claimed in any preceding claim containing a skin compatible mono or diester of glycerol with an aliphatic (C_{6-22}) carboxylic acid. | 5 |
| | 26. A composition as claimed in any preceeding claim containing a skin compatible ester having at least one hydroxyl group of a poly(2-10)glycerol with an aliphatic (C_{6-22}) carboxylic | |
| 10 | 27. A composition as claimed in any preceding claim containing a skin compatible mono- | 10 |
| | ether of a polyethylene-glycol with an aliphatic (C ₁₂₋₁₈) alcohol having an HLB value of from 10 to 18. 28. A composition as claimed in claim 27 wherein the mono ether is polyoxyethylene(10)o- | • |
| 15 | leyl ether. | 15 |
| 15 | 29. A composition as claimed in any preceding claim containing a skin compatible ester of an aliphatic (C_{6-22}) carboxylic acid with a) a polyethylene glycol | 13 |
| | b) a saccharose | |
| 20 | c) a sorbitan or d) a polyethylene glycol sorbitan ether, | 20 |
| | the ester having an HLB value of from 10 to 18. 30. A composition according to any preceeding claim containing as active agent (E)-N- | |
| | methyl-6, 6-dimethyl-N-(naphthylmethyl)hept-2-en-4-inyl-1-amine, naftifin, ketotifen, pizotifen, griseofulvin, fluocinolone acetonidie, triamcinolone acetonide, 14-0-[5-(2-amino-1,3,4-triazoly | 22 |
| 25 | thioacetyl]-dihydro-mutiline, or proquazone. 31. A composition according to any preceeding claim containing as active agent clemastine. | 25 |
| | 32. A composition according to any preceeding claim containing as active agent tizanidine. 33. A composition according to claim 30 containing 14-0-[5-(2-amino-1,3,4-triazolyl)thioa- | |
| 30 | cetyl]-dihydro-mutiline. 34. A composition according to claim 31 or 33 containing hexyl laurate, poly(7)ethylene | 30 |
| | glycol glyceryl cocoate and polyoxyethylene(10)oleyl ether. 35. A composition according to claim 32 containing 6 to 10% of tizanidine, | |
| | 15 to 25% of water-immisicible organic solvent, 15 to 25% of emulsifier, | |
| 35 | 5 to 10% of co-emulsifier, and 30 to 35% of water. | 35 |
| | 36. A composition according to claim 35 containing isopropyl laurate, polyoxyethylene(lo)oleyl ether and dodecanol. | |
| 40 | 37. A pharmaceutical composition in the form of a microemulsion, substantially as hereinbe- | 40 |
| 40 | fore described with reference to any one of the Examples. 38. A process for the production of a skin-penetrable pharmaceutical composition which | 40 |
| | comprises forming a microemulsion from water and a skin-penetrable pharmacologically active agent and skin compatible excipients capable of functioning as a water-immiscible organic solvent, an emulsifier and a co-emulsifier. | |
| 45 | | 45 |
| | form an emulsion and then cooled to form a microemulsion. | |
| | 40. A process for the production of a composition as defined in claim 1 substantially as hereinbefore described with reference to the Examples. | |
| 50 | 39 or 40. | 50 |
| | 42. A method of enhancing the penetration of a skin-penetrable pharmacologically active agent through the skin which comprises applying the active agent to the skin in the form of a | • |
| 55 | microemulsion consisting of skin compatible excipients. 43. A method according to claim 42 wherein the active agent is applied in the form of a | 55 |
| | microemulsion as defined in any one of claims 1 to 37. 44. Use of a microemulsion consisting of skin compatible excipients t administer percutane- | |
| | ously a skin-penetrable pharmacologically active agent. 45. Us according to claim 44 wh r in the active agent is tizanidine. | |
| 60 | 46. Us according to claim 44 wherein the active agent is clemastine. | 60 |
| | 47. A microemulsi n c mprising an active agent chosen from (E)-N-methyl-6,6-dimethyl-N-(naphthylmethyl)hept-2- n-4-inyl-1-amine, naftifin, ketotifen, piz tif n, griseofulvin, fluocinolane ac tonide, triamcinolone acetonide, 14-0-[5-(2-amino-1,3,4-triazolyl)thioacetyl]-dihydro-mutiline, | |
| 65 | r proquazone. 48. A microemulsion comprising clemastine r tizanidine. | 65 |
| | | |

- 49. A meth d of administering tizanidine by topical administration.
 50. A topical pharmaceutical composition comprising tizanidine.
- 51. A semi-solid pharmaceutical composition comprising tizanidine.

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